



09433360 Results

SEQ ID NO: 2

SUMMARIES

		8				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
		100 0	250	20	AAY57290	Human BGCKr protei
1 2	1819 1819	100.0	350 350	20	AAY17435	Human signal pepti
3	1819		350	20	AA117433 AAW93169	Human HFIAO41 prot
		100.0	350	21	AAY94325	Human seven transm
4	1819	100.0		21		Human CCR11 protei
5	1819	100.0	350		AAG80119	-
6	1819	100.0	350	22	AAU08994	Human G protein-co
7	1819	100.0	350	22	AAG67237	Amino acid sequenc Human chemokine re
8	1819	100.0	382	22	AAB62389	
9	1814	99.7	349	20	AAW93170	Human HFIAO41 prot
10	1814	99.7	350	20	AAY30125	A human seven-pass
11	1810	99.5	350	21	AAY71301	Human orphan G pro
12	1810	99.5	350	21	AAB02835	Human G protein co
13	1808	99.4	350	21	AAB37788	Human TSC7. Homo
14	1728	95.0	333	20	AAY57289	Human BGCKr partia
15	1620	89.1	350	22	AAG67238	Amino acid sequenc
16	1591	87.5	350	20	AAY57291	Mouse BGCKr protei
17	1275	70.1	246	20	AAY57292	Human BGCKr protei
18	1257	69.1	242	22	AAM99949	Human expressed po
19	1212	66.6	263	20	AAY30126	A seven-pass trans
20	862	47.4	164	22	ABB11162	Human orphan GPCR
21	862	47.4	164	22	AAM79310	Human protein SEQ
22	824	45.3	159	21	AAB41786	Human ORFX ORF1550
23	761	41.8	175	22	AAM99976	Human expressed po
24	761	41.8	175	22	ABB10276	Human cDNA SEQ ID
25	761	41.8	175	22	AAU18115	Novel human uterin
26	761	41.8	175	22	AAU18361	Human endocrine po
27	761	41.8	175	22	AAU18669	Renal and cardiova
28	761	41.8	175	22	AAU21655	Novel human neopla
29	659	36.2	358	15	AAR53745	Partial sequence o
30	659	36.2	358	21	AAB21689	Human 7TM receptor
31	659	36.2	378	19	AAW48724	Human V31 seven tr
32	659	36.2	378	21	AAB21688	Human 7TM receptor
33	659	36.2	378	22	AAG80114	Human CCR7 protein
34	659	36.2	378	22	AAB50859	Human CCR7. Homo
35	659	36.2	410	15	AAR53743	Putative seven tra
36	659	36.2	410	19	AAW48723	Polypeptide sequen
37	659	36.2	410	21	AAB21687	Genomic clone of 7
38	659	36.2	569	22	ABG12373	Novel human diagno
39	656	36.1	378	15	AAR53744	Putative seven tra
40	655	36.0	378	21	AAY90663	Human mutant G pro
41	653	35.9	378	21	AAY90629	Human G protein-co
42	650	35.7	378	21	AAB21699	7TM receptor prote
43	643	35.3	369	22	AAG80116	Human CCR9a protei
44	643	35.3	378	15	AAR54079	Epstein Barr virus
45	643	35.3	378	19	AAW56164	G-protein coupled
			2,0			- F

Issued:

Result		* Query				
No.	Score	Match	Length I	DB 	ID	Description
1	1819	100.0	350	2	US-08-966-316-16	Sequence 16, Appl
2	1620	89.1	350	2	US-08-966-316-18	Sequence 18, Appl
3	659	36.2	358	1	US-08-153-848-19	Sequence 19, Appl
4	659	36.2	358	3	US-09-299-843A-19	Sequence 19, Appl
5	659	36.2	358	4	US-09-088-337B-19	Sequence 19, Appl
6	659	36.2	358	5	PCT-US93-11153-19	Sequence 19, Appl

36.2 378 1 US-08-153-848-15 Sequence 15, Appl 659 378 3 US-09-299-843A-15 36.2 Sequence 15, Appl 8 659 36.2 378 4 US-09-251-545-1 Sequence 1, Appli 659 378 4 Sequence 15, Appl 10 US-09-088-337B-15 659 36.2 11 659 36.2 378 5 PCT-US93-11153-15 Sequence 15, Appl 410 1 US-08-153-848-7 12 659 36.2 Sequence 7, Appli 410 3 US-09-299-843A-7 13 659 36.2 Sequence 7, Appli 410 4 US-09-088-337B-7 410 5 PCT-US93-11153-7 Sequence 7, Appli Sequence 7, Appli 14 659 36.2 15 659 36.2 16 650 35.7 378 3 US-09-299-843A-66 Sequence 66, Appl 378 4 US-09-088-337B-66 650 17 35.7 Sequence 66, Appl 18 643 35.3 378 1 US-08-383-750-2 Sequence 2, Appli 378 1 US-08-383-751A-2 19 643 35.3 Sequence 2, Appli 20 643 35.3 378 3 US-08-352-678-2 Sequence 2, Appli 378 4 US-09-045-583-49 378 5 PCT-US93-09636-2 21 643 35.3 Sequence 49, Appl 22 643 35.3 Sequence 2, Appli 23 637 35.0 357 4 US-09-266-464-2 Sequence 2, Appli 359 1 US-08-153-848-24 24 635.5 34.9 Sequence 24, Appl Sequence 24, Appl 25 635.5 34.9 359 3 US-09-299-843A-24 359 4 US-09-088-337B-24 26 635.5 34.9 Sequence 24, Appl 27 635.5 34.9 359 5 PCT-US93-11153-24 Sequence 24, Appl 361 2 US-08-902-294-2 361 3 US-09-178-637-2 28 Sequence 2, Appli 606.5 33.3 29 606.5 33.3 Sequence 2, Appli 30 374 4 US-09-045-583-48 Sequence 48, Appl 583 32.1 31 581 31.9 342 4 US-09-116-498-4 Sequence 4, Appli Sequence 6, Appli 32 569 31.3 342 4 US-09-116-498-6 342 2 US-08-742-011-2 33 560 30.8 Sequence 2, Appli 342 4 US-09-275-384B-5 34 560 30.8 Sequence 5, Appli 35 560 30.8 342 4 US-09-116-498-2 Sequence 2, Appli 560 30.8 342 4 US-09-449-437A-2 36 Sequence 2, Appli 37 537.5 360 4 US-08-875-573-20 29.5 Sequence 20, Appl 38 537.5 29.5 360 4 US-09-232-878-2 Sequence 2, Appli 39 537.5 29.5 360 4 US-09-045-583-55 Sequence 55, Appl 355 1 US-07-759-568-1 40 534.5 29.4 Sequence 1, Appli 41 534.5 29.4 355 1 US-08-450-393A-8 Sequence 8, Appli 534.5 355 2 US-08-390-000A-5 42 29.4 Sequence 5, Appli 43 4 534.5 29.4 355 US-08-446-669-8 Sequence 8, Appli 355 5 PCT-US95-00476-8 44 534.5 29.4 Sequence 8, Appli 45 534.5 29.4 360 1 US-08-202-056-7 Sequence 7, Appli

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Sdo com & B
RESULT 1
US-08-966-316-16
; Sequence 16, Application US/08966316
; Patent No. 5932445
  GENERAL INFORMATION:
    APPLICANT: Lal, Preeti
    APPLICANT: Au-Young, Janice
    APPLICANT: Reddy, Roopa
APPLICANT: Murry, Lynn E.
    APPLICANT: Mathur, Preete
    TITLE OF INVENTION: SIGNAL PEPTIDE - CONTAINING PROTEINS
    NUMBER OF SEQUENCES: 18
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
      STREET: 3174 Porter Drive
      CITY: Palo Alto
      STATE: CA
      COUNTRY: USA
      ZIP: 94304
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ for Windows Version 2.0
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/966,316
      FILING DATE: Herewith
      CLASSIFICATION: 435
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PRIOR APPLICATION DATA:
      APPLICATION NUMBER:
      FILING DATE:
    ATTORNEY/AGENT INFORMATION:
     NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0424 US
    TELECOMMUNICATION INFORMATION:
     TELEPHONE: 650-855-0555
      TELEFAX: 650-845-4166
      TELEX:
  INFORMATION FOR SEQ ID NO: 16:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 350 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    IMMEDIATE SOURCE:
      LIBRARY: UTRSNOT11
      CLONE: 2547002
US-08-966-316-16
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US-08-966-316-18
; Sequence 18, Application US/08966316
; Patent No. 5932445
 GENERAL INFORMATION:
    APPLICANT: Lal, Preeti
    APPLICANT: Au-Young, Janice APPLICANT: Reddy, Roopa
    APPLICANT: Murry, Lynn E.
    APPLICANT: Mathur, Preete
    TITLE OF INVENTION: SIGNAL PEPTIDE - CONTAINING PROTEINS
    NUMBER OF SEQUENCES: 18
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Incyte Pharmaceuticals, Inc.
     STREET: 3174 Porter Drive
     CITY: Palo Alto
     STATE: CA
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COUNTRY: USA
      ZIP: 94304
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ for Windows Version 2.0
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/966,316
      FILING DATE: Herewith
      CLASSIFICATION: 435
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER:
      FILING DATE:
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0424 US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 650-855-0555
      TELEFAX: 650-845-4166
      TELEX:
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      TYPE: amino acid
     STRANDEDNESS: single
     TOPOLOGY: linear
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      CLONE: 399711
US-08-966-316-18
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        Db
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                              SUMMARIES
Result
             Ouerv
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No. Score Match Length DB ID Description



1620 89.1 350 2 JN0621 G protein-coupled 1 378 2 B55735 36.2 lymphocyte-specifi 2 659 650 35.7 378 2 A55735 G protein-coupled 3 643 35.3 378 2 A45680 G protein-coupled 4 5 583 32.1 369 2 JC5068 G protein-coupled 6 537.5 29.5 360 2 A57160 chemokine (C-C) re 7 534.5 29.4 360 2 A53611 interleukin-8 rece 8 524 28.8 354 I58186 probable G protein 9 522.5 28.7 355 2 JC4304 orphan G protein-c 10 522.5 28.7 360 2 JC4587 chemokine (C-C) re 518.5 354 2 B55733 11 28.5 G protein-coupled 12 511 28.1 358 2 A53752 interleukin-8 rece 501.5 27.6 355 2 J01231 interleukin-8 rece 13 14 500.5 27.5 352 2 A45747 neuropeptide Y/pep neuropeptide Y/pep 15 499.5 353 2 S28787 27.5 16 493.5 27.1 352 2 G00048 fusin (LESTRA) - c 17 492 27.0 360 2 JC2443 chemokine (C-C) re 18 489.5 350 2 A39445 interleukin-8 rece 26.9 19 486 26.7 355 2 JC5067 G protein-coupled 2 20 485 26.7 355 T49339 macrophage inflamm 21 484.5 26.6 352 2 A43113 chemokine (C-C) re 22 483.5 359 2 A48921 interleukin-8 rece 26.6 23 374 2 I38450 chemokine (C-C) re 482 26.5 479.5 359 2 149341 MIP-1 alpha recept 24 26.4 25 476.5 26.2 367 2 JE0349 interferon-inducib 26 474 26.1 355 2 A45177 chemokine (C-C) re 27 471 25.9 356 2 S42096 interleukin-8 rece 28 470 25.8 383 2 S55594 G protein-coupled 29 461.5 355 2 G02436 chemokine (C-C) re 25.4 30 2 455.5 25.0 359 S44425 angiotensin II rec 452.5 359 2 JC1104 31 24.9 angiotensin II rec 32 452.5 359 2 JC2134 24.9 angiotensin II rec 33 450 359 2 S15403 angiotensin II rec 24.7 34 450 24.7 374 2 S32785 G protein-coupled 35 446.5 24.5 359 2 A48857 angiotensin II rec 36 445.5 24.5 356 2 149340 MIP-1 alpha recept 37 442.5 24.3 372 2 S26667 G protein-coupled 38 442.5 24.3 374 2 S42628 G protein-coupled 39 362 2 JN0694 441 24.2 angiotensin II rec 40 439 24.1 359 2 A42656 angiotensin II rec 41 438.5 24.1 359 2 JH0621 angiotensin II rec 438.5 362 2 A39714 G protein-coupled 42 24.1 43 437.5 24.1 327 2 \$56162 MDCR15 protein - h 44 434 23.9 359 2 JQ1516 angiotensin II rec 359 2 I39418 45 428.5 23.6 angiotensin II rec

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JN0621
G protein-coupled receptor type B - bovine
C; Species: Bos primigenius taurus (cattle)
C;Date: 24-Feb-1994 #sequence revision 24-Feb-1994 #text change 19-May-2000
C; Accession: JN0621
R; Matsuoka, I.; Mori, T.; Aoki, J.; Sato, T.; Kurihara, K.
Biochem. Biophys. Res. Commun. 194, 504-511, 1993
A; Title: Identification of novel members of G-protein coupled receptor superfamily
expressed in bovine taste tissue.
A; Reference number: JN0621; MUID: 93326166
A; Accession: JN0621
A; Molecule type: mRNA
A; Residues: 1-350 < MAT>
A; Cross-references: GB:S63848; NID:g399710; PIDN:AAB27547.1; PID:g399711
A; Experimental source: tongue taste papillae
C; Comment: This protein is involved in modulating taste sensitivity or regeneration of
taste cells.
C; Superfamily: vertebrate rhodopsin
C; Keywords: G protein-coupled receptor; glycoprotein; receptor; transmembrane protein
F;42-66/Domain: transmembrane #status predicted <TM1>
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F;80-99/Domain: transmembrane #status predicted <TM2> F;114-135/Domain: transmembrane #status predicted <TM3> F;154-175/Domain: transmembrane #status predicted <TM4> F;200-222/Domain: transmembrane #status predicted <TM5> F;242-265/Domain: transmembrane #status predicted <TM6> F;284-306/Domain: transmembrane #status predicted <TM7> F;6,19/Binding site: carbohydrate (Asn) (covalent) #status predicted Query Match 89.1%; Score 1620; DB 2; Length 350; Best Local Similarity 86.0%; Pred. No. 1.1e-131; Matches 301; Conservative 27; Mismatches 22; Indels 0; Gaps ${\tt 1} {\tt MALEQNQSTDYYYEENEMNGTYDYSQYELICIKEDVREFAKVFLPVFLTIVFVIGLAGNS} {\tt 60} \\$ Qу 1 MAVEYNOSTDYYYEENEMNDTHDYSQYEVICIKEEVRKFAKVFLPAFFTIAFIIGLAGNS 60 Db 61 MVVAIYAYYKKQRTKTDVYILNLAVADLLLLFTLPFWAVNAVHGWVLGKIMCKITSALYT 120 Qy 61 TVVAIYAYYKKRTKTDVYILNLAVADLFLLFTLPFWAVNAVHGWVLGKIMCKVTSALYT 120 Db 121 LNFVSGMQFLACISIDRYVAVTKVPSQSGVGKPCWIICFCVWMAAILLSIPQLVFYTVND 180 Ov Db 121 VNFVSGMOFLACISTDRYWAVTKAPSOSGVGKPCWVICFCVWVAAILLSIPQLVFYTVNH 180 Qу 181 NARCIPIFPRYLGTSMKALIQMLEICIGFVVPFLIMGVCYFITARTLMKMPNIKISRPLK 240 Db 181 KARCVPIFPYHLGTSMKASIQILEICIGFIIPFLIMAVCYFITAKTLIKMPNIKKSQPLK 240 241 VLLTVVIVFIVTQLPYNIVKFCRAIDIIYSLITSCNMSKRMDIAIQVTESIALFHSCLNP 300 Qу 241 VLFTVVIVFIVTQLPYNIVKFCQAIDIIYSLITDCDMSKRMDVAIQITESIALFHSCLNP 300 Db 301 ILYVFMGASFKNYVMKVAKKYGSWRRQRQSVEEFPFDSEGPTEPTSTFSI 350 Qу Db 301 VLYVFMGTSFKNYIMKVAKKYGSWRRQRQNVEEIPFESEDATEPTSTFSI 350

Result No.	Score	Query Match	Length	DB	ID	Description
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3	659	36.2	378	1	CKR7_HUMAN	P32248 homo sapien
4	650	35.7	378	1	CKR7_MOUSE	P47774 mus musculu
5	639	35.1	369	1	CKR9_MOUSE	Q9wut7 mus musculu
6	637	35.0	357	1	CKR9_HUMAN	P51686 homo sapien
7	605	33.3	367	1	CKR6_MOUSE	O54689 mus musculu
8	583	32.1	374	1	CKR6_HUMAN	P51684 homo sapien
9	581	31.9	342	1	CCR6_CERAE	Ol8983 cercopithec
10	571	31.4	343	1	CCR6_MACMU	Q9xt45 macaca mula
11	569	31.3	342	1	CCR6_MACNE	O19024 macaca neme
12	560	30.8	342	1	CCR6_HUMAN	000574 homo sapien
13	537.5	29.5	360	1	CKR4_HUMAN	P51679 homo sapien
14	534.5	29.4	360	1	IL8B_HUMAN	P25025 homo sapien
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17	524	28.8	354	1	C3X1_RAT	P35411 rattus norv
18	523	28.8	384	1	CKD6_HUMAN	000590 homo sapien
19	522.5	28.7	353	1	IL8B_GORGO	Q28422 gorilla gor
20	522.5	28.7	355	1	C3X1_HUMAN	P49238 homo sapien
21	522.5	28.7	360	1	CKR4_MOUSE	P51680 mus musculu
22	521.5	28.7	353	1	IL8B_MACMU	Q28519 macaca mula
23	520.5	28.6	362	1	CKRA_MOUSE	Q9jl21 mus musculu
24	512	28.1	356	1	IL8B_CANFA	O97571 canis famil
25	511	28.1	354	1	C3X1_MOUSE	Q9z0d9 mus musculu
26	511	28.1	358	1	IL8B_RABIT	P35344 oryctolagus
27	504	27.7	360	1	CKR2_MACMU	O18793 macaca mula

355 1 IL8A_RABIT 28 501.5 27.6 P21109 oryctolagus 358 1 CKR3 CAVPO 501.5 27.6 Q9z2i3 cavia porce 29 352 1 CCR4_HUMAN 30 500.5 27.5 P30991 homo sapien 353 1 CCR4_BOVIN 360 1 IL8B_BOVIN 31 499.5 27.5 P25930 bos taurus Q28003 bos taurus 32 499.5 27.5 352 1 CCR4 PAPAN P56491 papio anubi 33 498.5 27.4 352 1 CKR5_CERTO 353 1 CCR4_FELCA 062743 cercocebus 498.5 34 27.4 35 498.5 27.4 P56498 felis silve 350 1 IL8A_GORGO 497.5 P55919 gorilla gor 27.4 36 37 495.5 27.2 352 1 CKR5_CERAE P56493 cercopithec 352 1 CCR4_CERTO 352 1 CCR4_MACFA 38 494.5 27.2 062747 cercocebus 39 493.5 27.1 Q28474 macaca fasc 352 1 CCR4 MACMU 40 493.5 27.1 P79394 macaca mula 359 1 CKR3_RAT O54814 rattus norv 41 492.5 27.1 359 1 IL8B_RAT 352 1 CKR5_PAPHA P35407 rattus norv 42 492.5 27.1 491.5 P56441 papio hamad 43 27.0 491.5 27.0 352 1 CKR5 PYGBI 097880 pygathrix b 490.5 27.0 352 1 CKR5 MACMU P79436 macaca mula 45

ALIGNMENTS

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DT
     16-OCT-2001 (Rel. 40, Last sequence update)
     16-OCT-2001 (Rel. 40, Last annotation update)
DT
DE
     C-C chemokine receptor type 11 (C-C CKR-11) (CC-CKR-11) (CCR-11)
DE
     (Chemokine receptor-like 1) (CCRL1) (CCX CKR).
     CCR11 OR CCBP2 OR VSHK1.
GN
     Homo sapiens (Human).
OS
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX
     NCBI TaxID=9606;
RN
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RP
     MEDLINE=20200450; PubMed=10734104;
RX
     Schweickart V.L., Epp A., Raport C.J., Gray P.W.;
RA
     "CCR11 Is a functional receptor for the monocyte chemoattractant
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     protein family of chemokines.";
     J. Biol. Chem. 275:9550-9556(2000).
RL
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     Gosling J., Dairaghi D.J., Wang Y., Hanley M., Talbot D., Miao Z.,
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RA
     Schall T.J.;
RT
     "Cutting edge: identification of a novel chemokine receptor that binds
     dendritic cell- and T cell-active chemokines including ELC, SLC, and
RT
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     TECK.":
RL
     J. Immunol. 164:2851-2856(2000).
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     Khoja H., Wang G., Ng C.-T.L., Tucker J., Brown T., Shyamala V.;
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     Gene 246:229-238(2000).
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CC
CC
         SCYA19/MIP3B/ELC, SCYA21/SLC AND SCYA25/TECK.
CC
     -!- SUBCELLULAR LOCATION: Integral membrane protein.
CC
     -!- TISSUE SPECIFICITY: PREDOMINANTLY EXPRESSED IN HEART. LOWER
CC
         EXPRESSION IN LUNG, PANCREAS, SPLEEN, SMALL INTESTINE AND FETAL
CC
         TISSUES.
CC
     -!- SIMILARITY: BELONGS TO FAMILY 1 OF G-PROTEIN COUPLED RECEPTORS.
CC
CC
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CC
     between the Swiss Institute of Bioinformatics and the EMBL outstation -
     the European Bioinformatics Institute. There are no restrictions on its
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    entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC
    or send an email to license@isb-sib.ch).
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DR
    EMBL; AF110640; AAF59827.1; -.
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DR
    PRINTS; PR00237; GPCRRHODOPSN.
DR
    PROSITE; PS00237; G PROTEIN RECEP F1 1; 1.
DR
    PROSITE; PS50262; G_PROTEIN_RECEP_F1_2; 1.
KW
    G-protein coupled receptor; Transmembrane; Glycoprotein.
FT
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    TRANSMEM
FT
                43
                      63
                              POTENTIAL.
FT
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FT
    TRANSMEM
                88
                     108
                              POTENTIAL.
FT
    DOMAIN
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                     113
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FΤ
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                     154
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FT
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               155
                     175
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                              CYTOPLASMIC (POTENTIAL).
FT
    DOMATN
               223
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FT
    TRANSMEM
               241
                     261
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                              EXTRACELLULAR (POTENTIAL).
FT
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                     289
FT
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               290
                     310
                              POTENTIAL.
FT
               311
    DOMAIN
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    CARBOHYD
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                      6
                              N-LINKED (GLCNAC. . .) (POTENTIAL) .
FT
    CARBOHYD
               19
                      19
                              N-LINKED (GLCNAC. . .) (POTENTIAL) .
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 Query Match 100.0%; Score 1819; DB 1; Length 350; Best Local Similarity 100.0%; Pred. No. 1.3e-104;
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 Matches 350; Conservative
                                             0; Indels
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Qу
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Qy
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Qy
     241 VLLTVVIVFIVTQLPYNIVKFCRAIDIIYSLITSCNMSKRMDIAIQVTESIALFHSCLNP 300
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CKRB_BOVIN
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                               PRT:
                                     350 AA.
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    P35350:
    01-JUN-1994 (Rel. 29, Created)
    01-JUN-1994 (Rel. 29, Last sequence update)
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16-OCT-2001 (Rel. 40, Last annotation update)
     C-C chemokine receptor type 11 (C-C CKR-11) (CC-CKR-11) (CCR-11)
DE
     (Possible gustatory receptor type B) (PPR1 protein).
GN
OS
     Bos taurus (Bovine).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
OC
OC
     Bovidae; Bovinae; Bos.
     NCBI_TaxID=9913;
OX
RN
     [1]
RР
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RC
     TISSUE=Tongue;
RX
     MEDLINE=93326166: PubMed=8392843:
     Matsuoka I., Mori T., Aoki J., Sato T., Kurihara K.;
RA
RT
     "Identification of novel members of G-protein coupled receptor
     superfamily expressed in bovine taste tissue.";
RT
RL
     Biochem. Biophys. Res. Commun. 194:504-511(1993).
CC
     -!- FUNCTION: RECEPTOR FOR SCYA2/MCP1, SCYA8/MCP2, SCYA13/MCP4.
CC
         SCYA19/MIP3B/ELC, SCYA21/SLC AND SCYA25/TECK (BY SIMILARITY).
CC
     -!- SUBCELLULAR LOCATION: Integral membrane protein.
CC
     -!- TISSUE SPECIFICITY: EXPRESSED IN CIRCUMVALLATE AND FUNGIFORM
CC
         PAPILLAE, OLFACTORY EPITHELIUM AND LUNG. LOWER EXPRESSION IN
CC
         LIVER, KIDNEY AND TONGUE EPITHELIUM BEARING NO TASTE PAPILLAE.
CC
         VERY LOW EXPRESSION IN THE CEREBRAL CORTEX OF THE BRAIN.
     -!- SIMILARITY: BELONGS TO FAMILY 1 OF G-PROTEIN COUPLED RECEPTORS.
CC
CC
CC
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     or send an email to license@isb-sib.ch).
CC
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     GCRDb; GCR 0757; -
     InterPro; IPR000276; GPCR Rhodpsn.
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     PROSITE; PS00237; G PROTEIN RECEP F1 1; 1.
DR
     PROSITE; PS50262; G_PROTEIN_RECEP_F1_2; 1.
KW
     G-protein coupled receptor; Transmembrane; Glycoprotein;
KW
     Phosphorylation.
FT
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                                  EXTRACELLULAR (POTENTIAL).
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FT
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FT
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FT
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FT
     CARBOHYD
                  6
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                                  N-LINKED (GLCNAC. . .) (POTENTIAL) .
FT
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FT
     DISULFID
                 112
                       184
                                 BY SIMILARITY.
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  Best Local Similarity 86.0%; Pred. No. 1.8e-92;
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Qy
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2	912	50.1	221	11	Q9ESK1	Q9esk1 rattus norv
3	643	35.3	369	4	Q9UQQ6	Q9uqq6 homo sapien
4	611	33.6	367	11	Q9R1V0	Q9r1v0 mus musculu
5	586.5	32.2	368	13	042444	042444 oncorhynchu
6	578	31.8	343	6	Q9N0Z0	Q9n0z0 cercocebus
7	566.5	31.1	351	11	Q9EQ16	Q9eq16 mus musculu
8	566	31.1	343	6	Q9BDS6	Q9bds6 macaca fasc
9	561.5	30.9	351	11	Q9ERH5	Q9erh5 mus musculu
10	560	30.8	342	4	Q9HCA5	Q9hca5 homo sapien
11	559	30.7	342	6	Q9TV16	Q9tv16 pan troglod
12	544	29.9	358	13	Q9PUA0	Q9pua0 acipenser r
13	534.5	29.4	360	11	Q91ZH4	Q91zh4 rattus norv
14	527	29.0	384	4	Q96A02	Q96a02 homo sapien
15	520.5	28.6	358	13	Q9YGC3	Q9ygc3 xenopus lae
16	516	28.4	355	11	Q9JLY8	Q9jly8 rattus norv
17	507.5	27.9	347	6	Q9MZM1	Q9mzm1 eulemur mac
18	507	27.9	357	13	042445	042445 oncorhynchu
19	506.5	27.8	347	6	Q9MZM7	Q9mzm7 callimico g
20	506.5	27.8	347	6	Q9MZM6	Q9mzm6 callicebus
21	503.5	27.7	347	6	Q9MZN0	Q9mzn0 alouatta se
22	503.5	27.7	352	6	Q9TV44	Q9tv44 cercopithec
23	502.5	27.6	347	6	Q9MZP1	Q9mzp1 mandrillus
24	502.5	27.6	347	6	Q9MZM9	Q9mzm9 ateles pani
25	502.5	27.6	347	6	Q9MZM8	Q9mzm8 callithrix
26	502.5	27.6	347	6	Q9MZM5	Q9mzm5 pithecia pi
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28	500.5	27.5	347	6	Q9MZN6	Q9mzn6 hylobates c
29	500.5	27.5	347	6	Q9MZN5	Q9mzn5 hylobates h
30	500.5	27.5	347	6	Q9MZN2	Q9mzn2 pongo pygma
31	500.5	27.5	347	6	Q9MZN1	Q9mzn1 gorilla gor
32	500.5	27.5	352	6	Q9TSQ8	Q9tsq8 cercopithec
33	499.5	27.5	339	6	Q9TUR8	Q9tur8 cercopithec
34	499.5	27.5	339	6	Q9TUR4	Q9tur4 mandrillus
35	499.5	27.5	347	6	Q9MZQ3	Q9mzq3 pygathrix a
36	499.5	27.5	347	6	Q9MZQ2	Q9mzq2 rhinopithec
37	499.5	27.5	347	6	Q9MZQ1	Q9mzq1 pygathrix b
38	499.5	27.5	347	6	Q9MZQ0	Q9mzq0 pygathrix n
39	499.5	27.5	347	6	Q9MZP9	Q9mzp9 nasalis lar
40	499.5	27.5	347	6	Q9MZP8	Q9mzp8 colobus pol
41 42	499.5	27.5	347	6 6	Q9MZP5	Q9mzp5 presbytis p
4.2	499.5	27.5	347	ь	Q9MZP4	Q9mzp4 presbytis f

43 499.5 27.5 347 6 Q9MZP3 Q9mzp3 presbytis e Q9mzp2 mandrillus 27.5 347 6 O9MZP2 44 499.5 499.5 27.5 347 6 Q9MZP0 Q9mzp0 macaca assa

ALIGNMENTS

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    01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
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OC
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OX
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    [1]
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RC
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RX
    MEDLINE=20519697; PubMed=11063828;
RA
    Dorf M.E., Berman M.A., Tanabe S., Heesen M., Luo Y.;
RT
    "Astrocytes express functional chemokine receptors.";
    J. Neuroimmunol. 111:109-121(2000).
RL
    EMBL; AF306532; AAK81712.1; -
DR
ĸw
    Receptor.
             350 AA; 39530 MW; C5F7D9DC949CECCF CRC64;
so
    SEQUENCE

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        Score 1596;
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        Length 350;

        Best Local Similarity
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 Matches 299; Conservative 26; Mismatches 25; Indels
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С	4	1825.8		162029	9	AC020632	AC020632 Homo sapi
-	5	1813.8	92.6	2407	6	E30057	E30057 G protein c
	6	1799.8	91.9	2156	6	E30058	E30058 G protein c
	7	1671.4	85.4	1925	6	AX244232	AX244232 Sequence
	8	1671.4	85.4	1925	9	AF193507	AF193507 Homo sapi
	9	1574.6	80.4	1839	6	AX244231	AX244231 Sequence
	10	1462.6	74.7	1637	6	AX041923	AX041923 Sequence
	11	1370.8	70.0	1388	6	AX244225	AX244225 Sequence
	12	1150	58.7	1150	6	AX244230	AX244230 Sequence
	13	1132	57.8	1147	6	AX113671	AX113671 Sequence
	14	1132	57.8	1147	6	AX203742	AX203742 Sequence
	15	1119.2	57.2	1147	6	AX113673	AX113673 Sequence
	16	1111.4	56.8	1113	6	AX239641	AX239641 Sequence
	17	1111.4	56.8	1113	9	AF233281	AF233281 Homo sapi
	18	1051.4	53.7	1053	6	AX239643	AX239643 Sequence
	19	1051.4	53.7	1053	6	AX244219	AX244219 Sequence
	20	961.2	49.1	2059	4	S63848	S63848 G-protein c
	21	961.2	49.1	2059	6	AX244221	AX244221 Sequence
	22	831	42.4	831	6	AX244223	AX244223 Sequence
	23	797	40.7	797	6	AX244229	AX244229 Sequence
	24	790.4	40.4	1138	10	AF306532	AF306532 Mus muscu
	25	604	30.8	604	6	AX244227	AX244227 Sequence
	26	481	24.6	481	6	AX244224	AX244224 Sequence
	27	447.4	22.8	665	10	AF090348	AF090348 Rattus no
	28	374	19.1	374	6	AX244228	AX244228 Sequence
	29	345.4	17.6	347	6	AX113681	AX113681 Sequence
	30	180.2	9.2	1631	10	AF121670	AF121670 Rattus no
	31	169.2	8.6	1302	10	AB009369	AB009369 Mus muscu
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	33	166.2	8.5	1500	9	HSDNABLR2	X84702 H.sapiens B
	34	166.2	8.5	1900	6	AR107241	AR107241 Sequence
	35	166.2	8.5	2058	6	AR107232	AR107232 Sequence
	36	166.2	8.5	2139	9	HUMEBI1CDN	L31581 Human G pro
	37	166.2	8.5	2160	6	AR107239	AR107239 Sequence
	38	166.2	8.5	2215	9	HUMEBI103	L31584 Human G pro
С	39	166.2		133801	9	AC004585	AC004585 Homo sapi
	40	166	8.5	1410	10	AB016031	AB016031 Mus muscu
	41	164.8	8.4	1186	9	HSA132337	AJ132337 Homo sapi
	42	164.8	8.4	2462	9	AF145440	AF145440 Homo sapi
	43	164.8	8.4	2544	9	AF145439	AF145439 Homo sapi
	44	164.8	8.4	2577	9	HSU45982	U45982 Human G pro
С	45	164.8	8.4	176968	9	AC005669	AC005669 Homo sapi

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	1	1958	100.0	1958	21	AAA27006	Human cDNA encodin
	2	1813.8	92.6	2407	20	AAX22557	Human HFIAO41 cDNA
	3	1800	91.9	1800	24	AAS98075	Human DNA for pote
	4	1799.8	91.9	2156	20	AAX22558	Human HFIAO41 cDNA
	5	1671.4	85.4	1925	22	AAH77722	Genomic clone of h
	6	1636.8	83.6	1660	20	AAX61288	Human signal pepti
	7	1574.6	80.4	1839	22	AAH77721	Consensuse cDNA se
	8	1462.6	74.7	1637	21	AAC68722	Human TSC7 cDNA.
	9	1370.8	70.0	1388	22	AAH77715	Nucleotide sequenc
:	10	1207.4	61.7	1232	20	AAZ90528	Human BGCKr protei
:	11	1150	58.7	1150	22	AAH77720	Nucleotide sequenc
:	12	1145.8	58.5	1150	20	AAX86674	DNA encoding a sev
:	13	1132	57.8	1147	22	AAF57685	Human chemokine re
:	14	1119.2	57.2	1147	22	AAF57686	Human chemokine re
:	15	1111.4	56.8	1113	22	AAS14572	Human cDNA encodin
:	16	1090	55.7	1130	20	AAZ90527	Human BGCKr partia
C	17	1057.4	54.0	1059	24	AAS98091	Human DNA for pote
:	18	1051.4	53.7	1053	22	AAH77711	Nucleotide sequenc
:	19	1041.8	53.2	1053	21	AAD01128	Human orphan G pro
	20	1041.8	53.2	1053	21	AAA46029	Human G protein co





	21	996.8	50.9	1001	21	AAC75995	Human ORFX ORF1550
	22	961.2	49.1	2059	22	AAH77712	Nucleotide sequenc
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	24	816	41.7	828	22	AAI99561	Human expressed po
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ALIGNMENTS

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; Sequence 17, Application US/08966316
; Patent No. 5932445
; GENERAL INFORMATION:
    APPLICANT: Lal, Preeti
    APPLICANT: Au-Young, Janice
    APPLICANT: Reddy, Roopa
APPLICANT: Murry, Lynn E
    APPLICANT: Mathur, Preete
     TITLE OF INVENTION: SIGNAL PEPTIDE - CONTAINING PROTEINS
    NUMBER OF SEQUENCES: 18
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
      STREET: 3174 Porter Drive
      CITY: Palo Alto
      STATE: CA
      COUNTRY: USA
      ZIP: 94304
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ for Windows Version 2.0
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/966,316
      FILING DATE: Herewith
      CLASSIFICATION: 435
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER:
      FILING DATE:
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0424 US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 650-855-0555
      TELEFAX: 650-845-4166
      TELEX:
  INFORMATION FOR SEQ ID NO: 17:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 1660 base pairs
      TYPE: nucleic acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    IMMEDIATE SOURCE:
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; GENERAL INFORMATION:
    APPLICANT: Lal, Preeti
APPLICANT: Au-Young, Janice
     APPLICANT: Reddy, Roopa
    APPLICANT: Murry, Lynn E. APPLICANT: Mathur, Preete
     TITLE OF INVENTION: SIGNAL PEPTIDE - CONTAINING PROTEINS
     NUMBER OF SEQUENCES: 18
     CORRESPONDENCE ADDRESS:
       ADDRESSEE: Incyte Pharmaceuticals, Inc.
       STREET: 3174 Porter Drive
       CITY: Palo Alto
STATE: CA
      COUNTRY: USA
      ZIP: 94304
     COMPUTER READABLE FORM:
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       COMPUTER: IBM Compatible
       OPERATING SYSTEM: DOS
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       FILING DATE: Herewith
      CLASSIFICATION: 435
     PRIOR APPLICATION DATA:
      APPLICATION NUMBER:
       FILING DATE:
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
       REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0424 US
     TELECOMMUNICATION INFORMATION:
      TELEPHONE: 650-855-0555
       TELEFAX: 650-845-4166
      TELEX:
  INFORMATION FOR SEQ ID NO: 17:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 1660 base pairs
      TYPE: nucleic acid
      STRANDEDNESS: single
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     CLONE: 2547002
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  GENERAL INFORMATION:
    APPLICANT: Lal, Preeti
APPLICANT: Au-Young, Janice
APPLICANT: Reddy, Roopa
     APPLICANT: Murry, Lynn E.
     APPLICANT: Mathur, Preete
     TITLE OF INVENTION: SIGNAL PEPTIDE - CONTAINING PROTEINS
     NUMBER OF SEQUENCES: 18
     CORRESPONDENCE ADDRESS:
       ADDRESSEE: Incyte Pharmaceuticals, Inc.
       STREET: 3174 Porter Drive
       CITY: Palo Alto
STATE: CA
       COUNTRY: USA
       ZIP: 94304
     COMPUTER READABLE FORM:
       MEDIUM TYPE: Diskette
       COMPUTER: IBM Compatible
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     CURRENT APPLICATION DATA:
       APPLICATION NUMBER: US/08/966,316
       FILING DATE: Herewith
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     PRIOR APPLICATION DATA:
       APPLICATION NUMBER:
       FILING DATE:
     ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
       REGISTRATION NUMBER: 36,749
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    TELEPHONE: 650-855-0555
    TELEFAX: 650-845-4166
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  INFORMATION FOR SEQ ID NO: 17:
   SEQUENCE CHARACTERISTICS:
    LENGTH: 1660 base pairs
    TYPE: nucleic acid
    STRANDEDNESS: single
    TOPOLOGY: linear
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    LIBRARY: UTRSNOT11
    CLONE: 2547002
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       1012 GTGTGGAGGAGTTTCCTTTTGATTCTGAGGGTCCTACAGAGCCAACCAGTACTTTTAGCA 1071
Db
    0v
       Db
    541 aataaaacatctgcattattctgaaactcaaatctcagacgccgtggttgcaacttataa 600
Qу
       Db
   1132 AATAAAACATCTGCATTATTCTGAAACTCAAATCTCAGACGCCGTGGTTGCAACTTATAA 1191
Qу
    601 taaagaatgggttgggggaaggggagaaataaaagccaagaagaggaaacaagataata 660
       Db
   1192 TAAAGAATGGGTTGGGGGAAGGGGGAGAAATAAAAGCCAAGAAGAGAAAACAAGATAATA 1251
    661 aatgtacaaaacatgaaaattaaaatgaacaatataggaaaataattgtaacaggcataa 720
0ν
       Db
   1252 AATGTACAAAACATGAAAATTAAAATGAACAATATAGGAAAATAATTGTAACAGGCATAA 1311
Qу
    721 gtgaataacactctgctgtaacgaagaagagctttgtgggtgataattttgtatcttggtt 780
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Db	1312	GTGAATAACACTCTGCTGTAACGAAGAAGAGCTTTGTGGTGATAATTTTGTATCTTGGTT 1371
Qу	781	gcagtggtgcttatacaaatctacacaagtgataaaatgacacagaactatatacacaca 840
Db	1372	GCAGTGGTGCTTATACAAATCTACACAAGTGATAAAATGACAGAGAACTATATACACACA 1431
Qy	841	ttgtaccaatttcaatttcctggttttgacattatagtataattatgtaagatggaacca 900
Db	1432	TTGTACCAATTTCAATTTCCTGGTTTTGACATTATAGTATAATTATGTAAGATGGAACCA 1491
Qу	901	ttggggaaaactgggtgaagggtacccaggaccactctgtaccatctttgtaacttcctg 960
Db	1492	TTGGGGAAAACTGGGTGAAGGGTACCCAGGACCACTCTGTACCATCTTTGTAACTTCCTG 1551
Qу	961	tgaatttataataatttcaaaataaaacaagtt-aaaaaaaaacccactatgctataagt 1019
Db	1552	TGAATTTATAATATTCAAAATAAACAAGTTAAAAAAAAACCCACTATGCTATAAGT 1611
Qу	1020	taggccatctaaaacagattattaaagaggttcatgttaaaaggcat 1066
Db	1612	TAGGCCATCTAAAACAGATTATTAAAGAGGTTCATGTTAAAAGGCAT 1658

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